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NEWS	1			Web Page for STN Seminar Schedule - N. America
NEWS	2	AUG	10	Time limit for inactive STN sessions doubles to 40 minutes
NEWS	3	AUG	18	COMPENDEX indexing changed for the Corporate Source (CS) field
NEWS	4	AUG	24	ENCOMPLIT/ENCOMPLIT2 reloaded and enhanced
NEWS	5			CA/CAplus enhanced with legal status information for
	-			U.S. patents
NEWS	6	SEP	09	50 Millionth Unique Chemical Substance Recorded in CAS REGISTRY
NEWS	7	SEP	11	WPIDS, WPINDEX, and WPIX now include Japanese FTERM
NEWS	8	OCT	21	thesaurus
NEWS				Derwent World Patents Index Coverage of Indian and Taiwanese Content Expanded
NEWS	9	OCT	21	Derwent World Patents Index enhanced with human translated claims for Chinese Applications and
				Utility Models
NEWS	10	NOV	23	Addition of SCAN format to selected STN databases
NEWS				Annual Reload of IFI Databases
NEWS				FRFULL Content and Search Enhancements
NEWS				DGENE, USGENE, and PCTGEN: new percent identity
112110		DLC	0 -	feature for sorting BLAST answer sets
NEWS	1.4	DEC	0.2	Derwent World Patent Index: Japanese FI-TERM
MEMP	1.4	DEC	02	thesaurus added
NEWS	1.5	DEC	0.2	PCTGEN enhanced with patent family and legal status
MEMO	10	DEC	02	display data from INPADOCDB
NEWS	16	DEC	02	USGENE: Enhanced coverage of bibliographic and
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DICTIONARY FILE UPDATES: 11 DEC 2009 HIGHEST RN 1197154-31-6

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ring nodes:
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22-23 23-24
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STRUCTURE UPLOADED

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*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

Structure attributes must be viewed using STN Express guery preparation.

=> s 11 sss full FULL SEARCH INITIATED 16:20:13 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED -1651 TO ITERATE

100.0% PROCESSED 1651 ITERATIONS SEARCH TIME: 00.00.01

267 ANSWERS

TOTAL ENTRY SESSION

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196.92

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=> file caplus COST IN U.S. DOLLARS

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FILE COVERS 1907 - 13 Dec 2009 VOL 151 ISS 25 FILE LAST UPDATED: 11 Dec 2009 (20091211/ED) REVISED CLASS FIELDS (/NCL) LAST RELOADED: Oct 2009 USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Aug 2009

CAplus now includes complete International Patent Classification (IPC)

reclassification data for the third quarter of 2009.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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L3
         17643 L2
=> s 13 and "hyaluronic acid"
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              1 "HYALURONICS"
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         74545 CONJUGATES
        127449 CONJUGATE
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          7330 LINKERS
         33734 LINKER
                 (LINKER OR LINKERS)
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        126576 LINKAGE
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6 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB The invention provides major histocompatibility complex (MHC) multimers comprising: (a) one or more MHC class I or class II antigens; (b) one or more antigenic peptides from pathogenic organisms (such as Borrelia) capable of binding to the MHC antigens; (c) linker mols.; and (d) multimerization domains that bind to the MHC complex and linker, and synthetic and recombinant methods for producing said

MHC multimers. The invention relates that said MHC multimers contain labels that include dyes, enzymes and/or radioactive mols., and that the multimers may contain an addnl. mol. related to a biol. activity, such as T cell activation, antigen presentation and/or therapy. The invention also relates that the multimerization domains are different types of carrier or scaffold mols., and include small mols., polymers, streptavidin, IgG, cells, liposomes and/or beads. The invention also provides the amino acid sequences of antigen peptides from Borrelia proteins, such as flagellins, outer membrane proteins and heat-shock proteins. The invention further provides for the use of said MHC multimers in immunization of individuals against diseases, such as Lyme disease, birrekusis and recurring fever, and in the diagnosis of a disease, and/or in the detection of T cells specific for a particular antigen. Finally, the invention provides for various methods used in detecting T cells specific for particular antigens.

ACCESSION NUMBER: 2009:1082060 CAPLUS

DOCUMENT NUMBER: 151:334871

TITLE:

Major histocompatibility complex (MHC) multimers specific for antigenic peptides from pathogens (such as Borrelia), their compositions, production and use in immunization, diagnosis and in detection of specific T cells

INVENTOR(S): Brix, Liselotte; Pedersen, Henrik; Scholler, Jorgen PATENT ASSIGNEE(S): Dako Denmark A/S, Den.

SOURCE: PCT Int. Appl., 2053pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT	NO.			KIN	_	DATE			APPL							
WC	2009	1060	73		A2									20081230			
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		FI,	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,
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							US 2008-83481P										
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RL: BSU (Biological study, unclassified); DGN (Diagnostic use); MOA (Modifier or additive use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(MHC multimers specific for antigenic peptides from pathogens (such as Borrelia), their compns., production and use in immunization, diagnosis and in detection of specific T cells)

59-05-2 CAPLUS RN

CM L-Glutamic acid, N-[4-[[(2,4-diamino-6-

pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

Absolute stereochemistry.

L6 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB The present invention describes novel methods to generate MHC Or HLA multimers and methods to improve existing and new MHC multimers. The invention also describes improved methods for the use of MHC multimers in anal. of T-cells in samples 5 including diagnostic and prognostic methods. Furthermore the use of MHC multimers in therapy are described, e.g. anti-tumor and anti-virus therapy, including isolation of antigen specific T-cells capable of inactivation or elimination of undesirable target cells or isolation of specific T-cells capable of regulation of other immune cells.

ACCESSION NUMBER: 2009:24490 CAPLUS

DOCUMENT NUMBER: 150:142453

TITLE: MHC multimers and conjugates for use in

diagnosis, prognosis and therapy of cancer, infection,

immune and autoimmune disease

INVENTOR(S): Brix, Liselotte; Pedersen, Henrik; Jakobsen, Tina; Schoeller, Joergen; Lohse, Jesper; Brunstedt, Katja;

Jacobsen, Kivin

PATENT ASSIGNEE(S): Dako Denmark A/S, Den. SOURCE: PCT Int. Appl., 470pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PA	ATENT				KIND DATE				APPL			DATE					
WO	2009				A1 20090108												
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										DK 2	007-	975			A 2	0070	703
										US 2	007-	9295	81P		P 2	0070	703

US 2007-929582P P 20070703 US 2007-929583P 20070703 US 2007-929586P Р 20070703

59-05-2, Methotrexate

RL: ARU (Analytical role, unclassified); DGN (Diagnostic use); MOA (Modifier or additive use); THU (Therapeutic use); ANST (Analytical study); BIOL (Biological study); USES (Uses)

(MHC multimers and conjugates for use in diagnosis, prognosis and therapy of cancer, infection, immune and autoimmune disease) 59-05-2 CAPLUS

CN L-Glutamic acid, N-[4-[[(2,4-diamino-6-

pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

Absolute stereochemistry.

REFERENCE COUNT:

THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

1.6 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB There is provided an organic-inorg, composite material containing a single nanoparticle therein, which is prepared by individually dispersing hydrophilic inorg. nanoparticles having a uniform particle size and conjugating biodegradable polymers to the surface of the nanoparticle, and a method of preparing the same. More particularly, the preparation method of

the

present invention comprises the following steps: (1) preparing hydrophilic nanoparticles by conjugating organic substances having a thiol group and a hydrophilic amine group to the surface of a core or a core/shell inorg. nanoparticle protected with a surfactant through a metal-thiolate (M-S) bond between them; (2) adjusting the concentration of the hydrophilic nanoparticles prepared in step (1) to 2+10-6 M or less and treating them in a sonication bath to prepare individually dispersed nanoparticles in the form of a single particle; and (3) conjugating biopolymers to the nanoparticle individually dispersed in step (2) through the formation of an amide bond between them under treatment in a sonication bath. The organic-inorg. composite material of the present invention exhibits high efficient photoluminescence and photostability as well as excellent chemical stability, dispersibility in water, biocompatibility and targetibility. Thus, it can be effectively used as a raw material for bioimaging or film coating. In an example, a hydrophobic CdSe/CdS quantum-dot solution was mixed with HSCH2CH2NH2·HCl to give a precipitate which is then conjugated with polyethylene glycol monomethyl ether mono(succinimidyl succinate) to prepare a composite.

ACCESSION NUMBER: 2008:224261 CAPLUS

DOCUMENT NUMBER: 148:239986

TITLE: Single nanoparticle containing organic-inorganic composite material and method of preparing the same

Woo, Kyoungja; Koo, Dong Hyun INVENTOR(S): PATENT ASSIGNEE(S): Korea Institute of Science & Technology, S. Korea SOURCE:

U.S. Pat. Appl. Publ., 16 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent English

LANGUAGE:

PRI

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATE	NO.	KIND	DATE	APPLICATION NO.		DATE	
US 20	0080044657	A1	20080221	US 2006-642772		20061219	
US 76	501391	B2	20091013				
KR 20	008017149	A	20080226	KR 2006-78757		20060821	
KR 80	09366	B1	20080305				
IORITY A	APPLN. INFO.:			KR 2006-78757	A	20060821	
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ASS OTHER SOURCE(S): MARPAT 148:239986

59-05-2, Methotrexate IT

RL: RGT (Reagent); RACT (Reactant or reagent)

(targeting agent; manufacture of single nanoparticle-containing organic-inorg. composite materials using hydrophilic linkers and

conjugation) 59-05-2 CAPLUS

RN CN L-Glutamic acid, N-[4-[[(2,4-diamino-6-

pteridinyl)methyllmethylaminolbenzovll- (CA INDEX NAME)

Absolute stereochemistry.

1.6 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB The present invention encompasses IL-12p40 binding proteins, particularly antibodies that bind human interleukin-12 (hIL-12) and/or human IL-23 (hIL-23). Specifically, the invention relates to antibodies that are chimeric, CDR grafted and humanized antibodies. Preferred antibodies have high affinity for hIL-12 and/or hIL-23 and neutralize h IL-12 and/or hIL-23 activity in vitro and in vivo. An antibody of the invention can be a full-length antibody or an antigen-binding portion thereof. Method of making and method of using the antibodies of the invention are also provided. The antibodies, or antibody portions, of the invention are useful for detecting hIL-12 and/or hIL-23 and for inhibiting hIL-12 and/or hIL-23 activity, e.g., in a human subject suffering from a disorder in which hIL-12 and/or hIL-23 activity is detrimental.

ACCESSION NUMBER: 2007:33392 CAPLUS

DOCUMENT NUMBER: 146:141003

TITLE: Human interleukin 12 subunit p40-binding antibodies, fragments and conjugates in combination with

other therapeutic agents for treating IL-12-associated acute and chronic inflammatory disease INVENTOR(S): Lacy, Susan E.; Fung, Emma; Belk, Jonathan P.; Dixon,

Richard W.; Roguska, Michael; Hinton, Paul R.; Kumar, Shankar

PATENT ASSIGNEE(S): SOURCE:

Abbott Laboratories, USA PCT Int. Appl., 211pp. CODEN: PIXXD2

APPLICATION NO.

DATE

DOCUMENT TYPE:

Patent English

KIND DATE

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION: PATENT NO.

		LENI .				KIND DATE					LICA								
	WO		0056	80												20060629			
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PRIORITY APPLN. INFO.:											WO	2005	-6956 -US25	79P 584		P 2	0050		

59-05-2, Methotrexate

RL: BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(human interleukin 12 subunit p40-binding antibodies, fragments and conjugates in combination with other therapeutic agents for treating IL-12-associated acute and chronic inflammatory disease)

RN 59-05-2 CAPLUS

CN L-Glutamic acid, N-[4-[[(2,4-diamino-6-

pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

Absolute stereochemistry.

```
ANSWER 5 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN
1.6
AB
    Disclosed is a hvaluronic acid/methotrexate compound
    useful as a therapeutic agent for joint diseases. The hyaluronic
     acid/methotrexate compound useful for the treatment of joint
     diseases comprises hyaluronic acid and methotrexate
     bonded to a hydroxy group of the acid through a linker having a
     peptide chain comprising one to eight amino acids, the linker
     being bonded to the hyaluronic acid through a
     carbamate group. Thus, a methotrexate derivative
     [MTX(Et)-α-PhePhe-NH-C10H2003-NH2] was prepared and reacted with
     p-nitrophenylchloroformate. The obtained phenylcarbamate compound
     4,7,10-trioxa-13-[N-[N-[N-[4-[[(2,4-diamino-6-
     pteridinyl)methyl]methylamino]benzoyl]-a-(05-
    methylglutamyl)]phenylalanyl]phenylalanylamino]tridecanylamine
     [MTX(Et)-\alpha-Phe-NH-C10H20O3-NHCO-O-C6H4-NO2] was reacted with
     sodium hyaluronate to give a hyaluronic acid
     /methotrexate compound of the present invention. The compound showed
     excellent antiarthritic effect in rat.
ACCESSION NUMBER:
                        2005:1103818 CAPLUS
DOCUMENT NUMBER:
                        143:392980
TITLE:
                        Hvaluronic acid/methotrexate
                        compound
                         Ikeya, Hitoshi; Morikawa, Tadashi; Takahashi, Koichi;
INVENTOR(S):
                         Izutani, Noriyuki; Tamura, Tatsuya; Okamachi, Akira;
                         Ishizawa, Takenori; Sato, Haruhiko; Higuchi,
                        Yoshinobu; Kato, Tatsuya; Honma, Akie
PATENT ASSIGNEE(S):
                        Denki Kagaku Kogyo Kabushiki Kaisha, Japan; Chugai
                         Seivaku Kabushiki Kaisha
SOURCE:
                        PCT Int. Appl., 81 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        Japanese
FAMILY ACC. NUM. COUNT: 2
PATENT INFORMATION:
     PATENT NO.
                        KIND DATE
                                          APPLICATION NO.
                                                                 DATE
     WO 2005095464
                        A1 20051013
                                         WO 2005-JP6472
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             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM,
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             RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
            MR, NE, SN, TD, TG
     EP 1739097
                         A1
                                20070103
                                         EP 2005-727780
                                                                  20050401
         R: DE, ES, FR, GB, IT
     US 20090093414
                         A1
                               20090409
                                           US 2008-547158
                                                                  20080529
                                                               A 20040402
PRIORITY APPLN. INFO .:
                                           JP 2004-110423
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                                            WO 2005-JP6472
                                                               W 20050401
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OTHER SOURCE(S):
                        MARPAT 143:392980
    59-05-2DP, Methotrexate, reaction products with linker
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RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU

peptides and hyaluronate

(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(hyaluronic acid/methotrexate compds. including peptide linkers for treatment of joint disease)

RN 59-05-2 CAPLUS

CN L-Glutamic acid, N-[4-[[(2,4-diamino-6-

pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD
(2 CITINGS)

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB Disclosed is a hyaluronic acid/methotrexate compound

useful as a therapeutic agent for joint diseases. The hyaluronic acid/methotrexate compound useful as a therapeutic agent for joint diseases comprises hyaluronic acid and methotrexate bonded to a carboxy group of the acid through a linker having a peptide chain comprising one to eight amino acids. For example,

2-[N-[N-[4-[[(2,4-diamino-6-pteridinyl)methyl]methylamino]benzoyl]α-(05-methylglutamyl)]phenylalanyl]phenylalanylamino]ethylamine
(MIX-α-Phen-MhC2H4NH2) was prepared and reacted with sodium

hyaluronate to obtain a conjugate, to examine its effect in arthritis model rats.

ACCESSION NUMBER: 2005:1004784 CAPLUS

DOCUMENT NUMBER: 143:292584

TITLE: Hyaluronic acid/methotrexate

compound

Sato, Haruhiko; Higuchi, Yoshinobu

PATENT ASSIGNEE(S): Denki Kaqaku Koqvo Kabushiki Kaisha, Japan; Chuqai

Seiyaku Kabushiki Kaisha

SOURCE: PCT Int. Appl., 95 pp.

CODEN: PIXXD2 Patent

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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WO	2005	0852	94		A1		20050915		WO 2005-JP3739						20050304				
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PRIORITY APPLN. INFO.:
                                            JP 2004-62616
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMA'
OTHER SOURCE(S): MARPAT 143:292584

IT 59-05-2D, Methotrexate, conjugates with

hvaluronic acids with peptide linkers

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (hyaluronic acid/methotrexate compds. with peptide

linkers for treatment of joint disease, and preparation thereof) RN 59-05-2 CAPLUS

L-Glutamic acid, N-[4-[[(2,4-diamino-6-

pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT:

THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)

REFERENCE COUNT:

22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB The present invention relates to methods for. the production of monomeric cytotoxic drug/carrier conjugates (the "conjugates") with higher drug loading and substantially reduced low conjugate fraction (LCF). Cytotoxic drug derivative/antibody conjugates, compns. comprising the conjugates and uses of the conjugates are also described. Particularly, the invention relates to anti-CD22 antibody-monomeric calichemicin conjugates.

The invention also relates to the conjugates of the invention, to methods of purification of the conjugates, to pharmaceutical compns. comprising the conjugates, and to uses of the conjugates.

ACCESSION NUMBER: 2003:892567 CAPLUS

DOCUMENT NUMBER: 139:386334

TITLE: Production of monomeric calicheamicin derivative

cytotoxic drug/carrier conjugates
INVENTOR(S): Kunz, Arthur; Moran, Justin Keith; Rubino, Joseph

Thomas; Jain, Neera; Vidunas, Eugene Joseph; Simpson, John McLean; Robbins, Paul David; Merchant, Nishith; Dijoseph, John Francis; Ruppen, Mark Edward; Damle, Nitin Krishnaji; Popplewell, Andrew Georce; et al.

PATENT ASSIGNEE(S): Wyeth Holdings Corporation, USA

SOURCE: PCT Int. Appl., 186 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

MO 2003092623 A2 20031113 WO 2003-US13910 200 MO 2003092623 A3 20040318 BB, BG, BR, BY, BZ, CA, C CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, G GM, HR, HU, JD, LL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, I LS, LT, LU, LV, MA, MD, MG, MK, MM, MM, MX, MZ, NI, NO, N PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, T TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RN: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, MZ, AM, AM KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, E FT, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SE BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, MM, MR, MR, SN, T CA 2483552 A1 20031113 CA 2003-2483552 200 EP 1507556 A2 20050223 EP 2003-724432 200 EP 1507556 A2 20050223 EP 2013-724432 200 EP 1507556 A2 20050227 CP 3013-11, LU, NL, SE, MR IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SE, MR IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SE, MR CN 1665532 A 20050018 BR 2003-9868 200 EN 2003009868 A 20051018 BR 2003-9868	H, CN, E, GH, K, LR, Z, OM,
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AU 2003231293 AI 20031117 AU 2003-231293 200 EP 1507556 A2 20050223 EP 2003-724432 200 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, M IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, S JP 2005524700 T 20050818 JP 2004-50080 CN 1665532 A 20050907 CN 2003-815260 200 CN 100482277 C 20090429	D, TG
EP 1507556 A2 20050223 EP 2003-724432 200 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, M	
EP 1507556 A2 20050223 EP 2003-724432 200 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, M	30502
IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, S JP 2005524700 T 20050818 JP 2004-500808 200 CN 1665532 A 20050907 CN 2003-815260 200 CN 100482277 C 20090429	30502
JP 2005524700 T 20050818 JP 2004-500808 200 CN 1665532 A 20050907 CN 2003-815260 200 CN 100482277 C 20090429	C, PT,
JP 2005524700 T 20050818 JP 2004-500808 200 CN 1665532 A 20050907 CN 2003-815260 200 CN 100482277 C 20090429	K
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BR 2003009868 A 20051018 BR 2003-9868 200	
	30502
NO 2004004663 A 20050125 NO 2004-4663 200	
MX 2004010792 A 20050307 MX 2004-10792 200	
TN 2004KN01802 A 20060106 TN 2004-KN1802 200	41129
IN 2007KN01141 A 20080801 IN 2007-KN1141 200	70402
AU 2009202609 A1 20090716 AU 2009-202609 200	90626
RITY APPLN. INFO.: US 2002-377440P P 200	
AU 2003-231293 A3 200	
	30502
IN 2004-KN1802 A3 200	

IT 59-05-2, Methotrexate

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (production of monomeric calicheamicin derivative cytotoxic drug/carrier conjugates)

PR

CN L-Glutamic acid, N-[4-[[(2,4-diamino-6-pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

RN 59-05-2 CAPLUS

OS.CITING REF COUNT: 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD

(5 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2009 ACS on STN

AB The present invention provides protein conjugates having a

glucose-aminoglycan-targeting domain conjugated directly or indirectly to a therapeutically useful protein via chemical or peptidyl linkage.

A conjugate of the invention is disclosed in which a

hyaluronan-binding protein is a receptor for hyaluronic acid-mediated mobility (RHAMM). The protein conjugates

selectively target certain tissues and organs and are useful for treating or preventing various physiol. and pathol. conditions. Methods of their use and preparation are described.

ACCESSION NUMBER: 2001:798086 CAPLUS

DOCUMENT NUMBER: 2001:798086 CAPLUS

TITLE: RHAMM peptide conjugates for drug targeting

INVENTOR(S): Woloski, B. Michael R.; Williams, Ashley Martin; Sereda, Terrance Jimmy; Wiebe, Deanna June

PATENT ASSIGNEE(S): Cangene Corporation, Can. SOURCE: PCT Int. Appl., 121 pp.

SOURCE: PCT Int. Appl., 121 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT :	NO.			KIND DATE				APPLICATION NO.							DATE			
					A2 20011101 A3 20020906				WO 2	001-	CA53	20010420							
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	2406 1274	593			A1		2001	1101	-	CA 2	001-	2406	593		2				
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	US 20040037834 PRIORITY APPLN. INFO.:						2004	0226		US 2 WO 2	000-	1986	13P	1	P 2	0030 0000 0010	420		

OTHER SOURCE(S): MARPAT 135:348866

- IT 59-05-2DP, Methotrexate, RHAMM conjugates RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
- (RHAMM peptide conjugates for drug targeting)
 RN 59-05-2 CAPLUS
- CN L-Glutamic acid, N-[4-[[(2,4-diamino-6pteridinyl)methyl]methylamino]benzoyl]- (CA INDEX NAME)

Absolute stereochemistry.

OS.CITING REF COUNT: 7 THERE ARE 7 CAPLUS RECORDS THAT CITE THIS RECORD (7 CITINGS)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD, ALL CITATIONS AVAILABLE IN THE RE FORMAT

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